DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. 2004N-0050]

Over-the-Counter Drug Products; Safety and Efficacy Review; Additional Dandruff Control Ingredient

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice of eligibility; request for data and information.

SUMMARY: The Food and Drug Administration (FDA) is announcing a call-fordata for safety and effectiveness information on the following conditions as part of FDA's ongoing review of over-the-counter (OTC) drug products: Piroctone olamine, 0.05 percent to 0.5 percent and 0.1 percent to 1.0 percent, for use as a dandruff control single active ingredient in leave-on and rinse-off dosage forms, respectively. FDA has reviewed a time and extent application (TEA) for these conditions and determined that they are eligible for consideration in its OTC drug monograph system. FDA will evaluate the submitted data and information to determine whether these conditions can be generally recognized as safe and effective (GRAS/E) for their proposed OTC use.

DATES: Submit data, information, and general comments by [insert date 90 days after date of publication in the **Federal Register**].

ADDRESSES: Submit written comments, data, and information to the Division of Dockets Management (HFA–305), Food and Drug Administration, 5630 Fishers Lane, rm. 1061, Rockville, MD 20852. Submit electronic comments, data, and information to http://www.fda.gov/dockets/ecomments.

FOR FURTHER INFORMATION CONTACT: Michael L. Koenig, Center for Drug Evaluation and Research (HFD–560), Food and Drug Administration, 5600 Fishers Lane, Rockville, MD 20857, 301–827–2222.

SUPPLEMENTARY INFORMATION:

I. Background

In the Federal Register of January 23, 2002 (67 FR 3060), FDA published a final rule establishing criteria and procedures for additional conditions to become eligible for consideration in the OTC drug monograph system. These criteria and procedures, codified in § 330.14 (21 CFR 330.14), permit OTC drugs initially marketed in the United States after the OTC drug review began in 1972 and OTC drugs without any marketing experience in the United States to become eligible for FDA's OTC drug monograph system. The term "condition" means an active ingredient or botanical drug substance (or a combination of active ingredients or botanical drug substances), dosage form, dosage strength, or route of administration, marketed for a specific OTC use (§ 330.14(a)). The criteria and procedures also permit conditions that are regulated as cosmetics or dietary supplements in foreign countries but that would be regulated as OTC drugs in the United States to become eligible for the OTC drug monograph system.

Sponsors must provide specific data and information in a TEA to demonstrate that the condition has been marketed for a material time and to a material extent to become eligible for consideration in the OTC drug monograph system. When the condition is found eligible, FDA publishes a notice of eligibility and request for safety and effectiveness data for the proposed OTC use. The TEA that the agency reviewed (Ref. 1) and FDA's evaluation of the TEA (Ref. 2) have been placed on public display in the

Division of Dockets Management (see **ADDRESSES**) under the docket number found in brackets in the heading of this document. Information deemed confidential under 18 U.S.C. 1905, 5 U.S.C. 552(b), or 21 U.S.C. 331(j) was deleted from the TEA before it was placed on public display.

II. Request for Data and Information

The conditions piroctone olamine, 0.05 percent to 0.5 percent and 0.1 percent to 1.0 percent, as a dandruff control single active ingredient in leaveon and rinse-off dosage forms, respectively, will be evaluated for inclusion in the monograph for OTC drug products for the control of dandruff, seborrheic dermatitis, and psoriasis (21 CFR part 358, subpart H). Accordingly, FDA invites all interested persons to submit data and information, as described in § 330.14(f), on the safety and effectiveness of these conditions for FDA to determine whether they can be GRAS/E and not misbranded under recommended conditions of OTC use. The TEA did not include an official or proposed United States Pharmacopeia-National Formulary (USP-NF) drug monograph for piroctone olamine. According to § 330.14(i), an official or proposed USP-NF monograph for piroctone olamine must be included as part of the safety and effectiveness data for this ingredient. Interested parties should provide an official or proposed USP-NF monograph and safety and effectiveness data for both leave-on and rinse-off dosage forms containing this ingredient.

Interested persons should submit comments, data, and information to the Division of Dockets Management (see ADDRESSES) by [insert date 90 days after date of publication in the Federal Register]. Three copies of all comments, data, and information are to be submitted. Individuals submitting written information or anyone submitting electronic comments may submit one copy.

Submissions are to be identified with the docket number found in brackets in the heading of this document and may be accompanied by supporting information. Received submissions may be seen in the Division of Dockets Management between 9 a.m. and 4 p.m., Monday through Friday. Information submitted after the closing date will not be considered except by petition under 21 CFR 10.30.

III. Marketing Policy

Under § 330.14(h), any product containing the conditions for which data and information are requested may not be marketed as an OTC drug in the United States at this time unless it is the subject of an approved new drug application or abbreviated new drug application.

IV. References

The following references are on display in the Division of Dockets

Management (see ADDRESSES) and may be seen by interested persons between

9 a.m. and 4 p.m., Monday through Friday.

- 1. TEA for piroctone olamine submitted by Keller and Heckman LLP on behalf of Clariant Gmbh., dated July 11, 2003.
 - 2. FDA's evaluation and comments on the TEA for piroctone olamine.

Dated: February 10, 2004.

Jeffrey Shuren,

Assistant Commissioner for Policy.

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